



CHARACTERIZATION OF ACTIVATED CARBON USING CHEMICAL ACTIVATION VIA MICROWAVE ULTRASONIC SYSTEM

(Pencirian Karbon Teraktif Menggunakan Sistem Pengaktifan Kimia Melalui Ketuhar Gelombang Ultrasonik)

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Abstract

Higher adsorption capacities of activated carbon (AC) can be evaluate according to pore volume, porosity and surface area. AC with higher pore volume, porosity and surface area desired in the present study in order to enhance the properties of AC supercapacitor. Thus, the present studies focus on the chemical activation process to increase the characterization of AC. The study was using bamboo waste as a precursor and the activation process was conducted using microwave ultrasonic system. The chemical agent used during the process was KOH and H₂SO₄. Microwave activation was conducted at intensity 100 W and 300 W for 30 min and sonication frequency was constantly set at 200 Hz for 60 min. The sample was carbonized at temperature of 400 °C and 800 °C using furnace for 2 hours. Then, AC was characterized for surface area using BET analysis and functioning group using FTIR analysis. The results shown the carboxyl, aliphatic, aromatic and phenolic hydroxyl group are present on raw bamboo while new functional group such as alkyl halide and some of some weak bands appeared which analogous with out of plane bending mode of the C-H or O-H group occur for AC. Active surface area and total pore volume of AC supercapacitor in 5M of concentration for H₂SO₄ and KOH corresponded to 1167 m²/g, 0.724 cm³/g, 740.10 m²/g, 0.462 cm³/g, respectively.

Keywords: supercapacitor, microwave-ultrasonic, activation, chemical, carbonization

Abstrak

Kapasiti penjerapan yang tinggi pada karbon teraktif boleh dinilai mengikut isipadu liang, keliangan dan luas permukaan. Keaktifan karbon dengan isi padu liang yang tinggi, tahap keliangan yang besar dan luas permukaan yang tinggi amat dikehendaki di dalam kajian ini bagi meningkatkan sifat-sifat keaktifan karbon superkapasitor. Oleh itu, kajian ini memberi tumpuan kepada proses pengaktifan kimia untuk meningkatkan pencirian yang diperlukan dalam karbon aktif. Kajian ini telah menggunakan sisa buluh sebagai pelopor dan proses pengaktifan telah dijalankan menggunakan sistem ultrasonik-gelombang mikro. Agen kimia yang digunakan semasa proses pengaktifan adalah KOH dan H₂SO₄. Pengaktifan gelombang ketuhar telah digunakan pada intensiti 100 W dan 300 W selama 30 min dan kekerapan sonikasi telah ditetapkan pada 200 Hz selama 60 min. Sampel telah dikarbonisasi pada suhu 400 °C dan 800 °C menggunakan relau selama 2 jam. Kemudian, karbon aktif dikelaskan bagi mendapatkan luas permukaan dengan menggunakan analisis BET dan kumpulan berfungsi pula menggunakan analisis FTIR. Keputusan menunjukkan karboksil aliphatik, aromatik dan kumpulan fenolik hidrosil hadir pada bahan asas buluh manakala kumpulan berfungsi baharu seperti alkil halida dan beberapa kumpulan yang lemah muncul dimana serupa dengan lenturan mod bagi C-H atau O-H yang terhasil pada karbon aktif. Kawasan permukaan yang aktif dan jumlah isipadu liang

karbon aktif superkapasitor pada kepekatan 5M untuk H_2SO_4 and KOH sepadan mengikut aturan 1167 m^2/g , 0.724 cm^3/g , 740.10 m^2/g , 0.462 cm^3/g .

Kata kunci: superkapasitor, gelombang mikro-ultrasonik, pengaktifan, bahan kimia, karbonisasi

Introduction

Activated carbon (AC) is widely use as catalyst, catalyst support and adsorbent for the removal of pollutant order from liquid to gases and for purification or recovery of chemicals [1]. A wide range of carbonaceous materials such as agricultural wastes or industrial wastes, coconut shell, coal and wood can be used as activated carbon precursor. In industrial practice, coal and coconut shell are two major sources for the production of activated carbon [2]. Bamboo, one of the lignocellulosic resources is a potential precursor for activated carbon due to its flexible, efficient and economical materials [3]. Bamboo has a high volatile matter and low of ash content that required in AC processing with high adsorption capacities are related to property of pore volume, porosity and surface area. Current approach in AC processing gave AC with a small of surface area, limited power intensity and energy capacity and low of porosity [4]. In order to overcome these problems, chemical activation and microwave ultrasonic activation has been introduced to enlarge the surface area, extensive the power intensity and energy capacity and enhance the porosity of AC.

Activated carbon works by the process of adsorption. Adsorption is a process when one material adheres to the surface of another material by means of physical or chemical attraction between the materials. There are two processes involves in development the formation of AC which are activation and carbonization processes. In activation, there are two types of method in order to produce activated carbon from carbonaceous material which is chemical and physical method. Physical activation procedure required two step of process which are carbonization and followed by activation using steam, oxygen or carbon dioxide as an activation agent while chemical activation procedure involves a single stage by using chemical as activating agent such as zinc chloride, potassium hydroxide, phosphoric acid and others [5]. Chemical procedure takes a few of advantage than physical method which has higher carbon yield and better developed of pore structure [6].

Thus, in the present study, bamboo was prepared for AC by using different of chemical activation agent which is KOH and H_2SO_4 under microwave ultrasonic radiation for activation process and continues with a single step of carbonization process at different temperature. The aim of this study is to characterize the characteristics of bamboo properties as activated carbon with chemical activation in microwave-ultrasonic activation.

Materials and Methods

Materials

The chemicals used in this study are sulphuric acid (H_2SO_4) and potassium hydroxide (KOH) purchased from R&M Chemical, Inc. Both chemicals were analytical grade.

Preparation of activated carbon

The precursor for activated carbon was residual bamboo obtained from Forest Research Institute Malaysia (FRIM). A total 50 g of dry bamboo was placed into crucible and the chemical agent, KOH or H_2SO_4 was added into the sample with ratio 1:1. Then, the sample undergoes microwave activation with different energy power for 2 hr. Activate sample was cooled and washed with distilled water until pH 6 to 7. The sample was dried in oven at 110 °C for 24 hours and stored in desiccator before undergoes carbonization process. Next, the sample was carbonized in the Vermont Furnace (Carbolite, Keison) under the flow of 150 cm^3/min nitrogen for 2 hours. After carbonization process, the activated carbon was cooled to ambient temperature for further analysis. Parameter used in the present study is stated in Table 1.

Table 1. Parameter used in Microwave-ultrasonic system with chemical agent

Sample	Experiment Procedure	Chemical Activating Agent
AC_OH_8	Ultrasonic-MW (300W)-Carbonize (800°C)	Base - KOH
AC_OH_4	Ultrasonic-MW (300W)-Carbonize (400°C)	Base - KOH
AC_H_8	Ultrasonic-MW (300W)-Carbonize (800°C)	Acid - H ₂ SO ₄
AC_H_4	Ultrasonic-MW (300W)-Carbonize (400°C)	Acid - H ₂ SO ₄
AC_K_8	Ultrasonic-MW (100W)-Carbonize (800°C)	Base - KOH
AC_K_4	Ultrasonic-MW (100W)-Carbonize (400°C)	Base - KOH
AC_SO_8	Ultrasonic-MW (100W)-Carbonize (800°C)	Acid - H ₂ SO ₄
AC_SO_4	Ultrasonic-MW (100W)-Carbonize (400°C)	Acid - H ₂ SO ₄

Proximate analysis of precursor

The raw material sample was characterized for proximate analysis by using American Society for Testing and Materials standard in order to determine the moisture content of bamboo (ASTM D 2867-99), the presence of volatile matter (ASTM D 5832-98), the ashes content (ASTM D 2866-94) and the amount of fixed carbon and waxes composition [7].

Characterization of activated carbon

Brunauer, Emmett, Teller (BET) test (Xyrapcorp, Autosorb-1, Malaysia) was used in order to determine the specific surface area of a variety materials by the BET Nitrogen adsorption technique. The adsorption process handled in 1 hour by using quick single point and multipoint particular BET surface area determinations.

Field Emission Scanning Electron Microscope (FESEM) was carried out using Carl Zeiss, Supra 40VP, Germany in conducive to examine the morphology of the sample and to analyze the composition, distribution and phase structure of residual bamboo. The sample must be electrically connected to the sample holder in order to prevent charging and distortion of the image.

Fourier Transform Infrared Spectroscopy (FTIR) (Perkin Elmer, Spectrum One-FTIR, USA) was performed in order to identify chemical bonds in a molecule by producing an infrared absorption spectrum for functional group classification. FTIR creates the absorbance spectra representing the unique chemical bonds and the molecular structure of the sample material. FTIR spectra were recorded between 4000 and 500 cm⁻¹ by using AVATAR 360 Spectrophotometer.

Results and Discussion

Determination of surface area and pore size

Table 2 shows the total of active surface area and total pore volume analyzed using nitrogen adsorption/desorption. The acquired results determined the surface of the chemical activation with base, KOH consist a greater surface area as well as pore volume as compared with chemical activation using acid activating agent, H₂SO₄. The carbonization of bamboo at 800 °C gave a higher active surface area then carbonization at 400 °C. The highest of active surface area obtained was 1162.73 m²/g with total pore value 0.7237 cm³/g using 300W of MW-ultrasonic radiation power. The active surface area obtain from the study was compare with previous research on AC derived from different agricultural waste and tabulated in Table 3. The result indicated that present study gave highest V_{mic} and active surface area is at the power with another AC using KOH as activating agent. This comparison showed bamboo is one of the efficient and effective precursor in produce a high of V_{mic} carbon.

Table 2. Chemical properties of produced activated carbon

Experiment	Active Surface Area (m ² /g)	Total Pore Volume (cm ³ /g)
AC_OH_8	1162.73	0.7237
AC_OH_4	2.8791	0.0024
AC_H_8	740.09	0.4019
AC_H_4	478.14	0.2262
AC_K_8	24.6062	0.0172
AC_K_4	1.8299	0.0063
AC_SO_8	-	0.0066
AC_SO_4	4.7868	0.0078

Table 3. Pore structure from other agricultural wastes by MW with KOH activation

Precursor	Power (W)	S _{BET} (m ² /g)	V _{mic} (cm ³ /g)	Ref
Bamboo	300	1162.73	0.7237	This work
Oil palm fiber	600	1223.00	0.4200	[8]
Coconut husk	600	1356.30	0.3920	[9]
Pineapple	600	1006.00	0.2800	[10]
Siris seed pods	620	1824.80	0.6450	[11]

Proximate analysis

Table 4 show the properties of raw bamboo precursor. Before undergoes the proximate analysis, raw bamboo has been dried into oven for 30 min (BamP) and 15 min (BamPr) of temperature in order to remove impurities and water content. From the results, it showed that BamP consist of low moisture content when compared to BamPr. For volatile and ash content, sample BamP gave a higher value which is 45.320% and 1.98% while BamPr sample obtain 40.178% of volatile content and 1.5% of ash content. For fixed carbon content BamPr sample have 58.322% whereas BamP consist of 52.7% of fixed carbon. High ash content is undesired in AC processing because it represents amount of inorganic material from the precursor activation and a numerous treatment needed to be take place [12].

Table 4. Proximate of raw bamboo precursor

Experiment	Moisture Content (%)	Volatile Content (%)	Ash Content (%)	Fixed Carbon (%)
BamP	13.300	45.320	1.98	52.7
BamPr	14.356	40.178	1.5	58.322

Elemental analysis (CHNS-O analysis)

Table 5 summarizes the results of elemental analysis which gave the percentage of carbon, hydrogen, nitrogen and sulphur present in the sample. The result indicated that the largest elements inside activated carbon are oxygen and carbon. With increment of activation temperature, there was decreasing in carbon content which probably due to the discharge of volatiles [13]. In chemical activation of acid activating agent, H₂SO₄ the activation process has

provoked a progress of discharge oxygen with notable loss at 800 °C. Discharge of oxygen happened because element of oxygen intimate the fraction of oxygen content was not located around the pore entry of activated carbon [14]. High of moisture and volatile content also give a higher value for oxygen element. On the contrary, there was a decreasing of carbon content in a higher temperature. Besides that, there was a reduction of hydrogen element at a temperature 800 °C. The percentages of carbon significantly high at temperature 400 °C with radiation power 300 W and 100 W based on different type of activating agent. The precursor of carbon could barely be heated without impregnation of activation agent, demonstrating that activation agent acted as the fundamental MW absorber at the primary stage. With the evolution of pore structure, the AC itself could acquire MW energy [15]. At low microwave energy, the pore structure was not satisfactorily created which improved with the expanding of microwave power. Nevertheless, at tremendous radiation power, absorbed MW energy eclipse at some of power so that the overfull energy keep create a small feature of carbon burnt and the structure was destroyed [16].

Table 5. Result of elemental analysis

Experiment	Carbon (%)	Hydrogen (%)	Nitrogen (%)	Sulphur (%)	Oxygen (%)
AC_OH_4	30.3338	2.5647	0.4022	0.0000	66.6993
AC_H_4	37.5033	0.8403	0.2574	0.0386	61.3604
AC_OH_8	20.5301	1.8585	1.1981	0.0000	76.4133
AC_H_8	14.0534	0.1058	0.399	0.0167	85.4251
AC_K_4	28.5147	2.4452	0.380	0.0130	68.6471
AC_SO_4	26.2542	1.8933	0.149	0.0000	71.7035
AC_K_8	21.4699	0.6440	0.412	0.0471	77.4270
AC_SO_8	20.7577	0.7182	0.466	0.0000	78.0581

FTIR analysis

The prepared activated carbon was analyzed for functional group using FTIR shown a following bands: 3308.78 cm^{-1} was attributed to O-H vibration in hydroxyl groups. The location of hydrogen bonded OH groups usually at range 3200 – 3750 cm^{-1} for alcohol and phenol which involve in hydrogen bonding may be due to adsorbed water [17], C=C stretching vibration of aromatic rings at 1599.98 cm^{-1} , the relatively intense band at 1031.89 cm^{-1} can be assigned to alcohol or phenol groups of R-OH, 1234.05 cm^{-1} with absence of C=O stretching from the esters (C-O) of strong intensity absorptions. Figure 1 shown band at range 756 – 540 cm^{-1} with 667.07 cm^{-1} is (C-O-H) twist abroad with strong aliphatic, 1350.48 cm^{-1} of amines compound occurs which correlated out of plane bending absorption, alkanes (C-H CH_2) found at range of 1457.03 cm^{-1} bending absorption of methylene groups and 1784.93 cm^{-1} denotes C=O stretching from ketones (C=OR), aldehyde (C=OH) and carboxylic acid (COOH). Some weak bands also appeared in the range of 600 – 900 cm^{-1} which associated with out-of-plane bending mode of (C-H) and O-H group. Heating at a high of temperature make some of peaks disappear in AC_K_8 and AC_H_8. The main surface functional groups present on the activated carbon of bamboo are lactones, ketones, carboxylic anhydrides, quinine structure and aromatic ring [18].

Activated carbon morphology

Figure 2 indicates the FESEM images of the raw bamboo and derived AC. There was no pore development on the precursor of raw bamboo residual and the structure of the surface is rough and uneven. From Figure 1a, many large pores shape and size were clearly found on the AC surface after the activation process of MW-ultrasonic and chemical activation. The well-developed pores had led to the large surface area and porous structure of the activated carbon had shown in Figure 1c and 1d. It can be seen that there was a significant difference between sample from KOH and H_2SO_4 surface texture. Pore development was caused by the breakdown of some material in the precursor due to thermal expansion during the activation step [19]. The reaction rate between the activating agent which is KOH and carbon also increase when the precursor is subjected to high activation temperature, thus leading to the formation of well-developed pores [20].

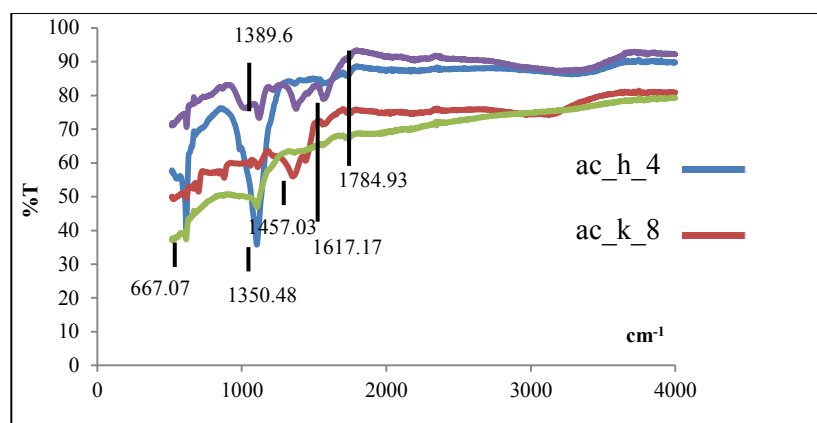


Figure 1. FTIR Spectra of activated carbon

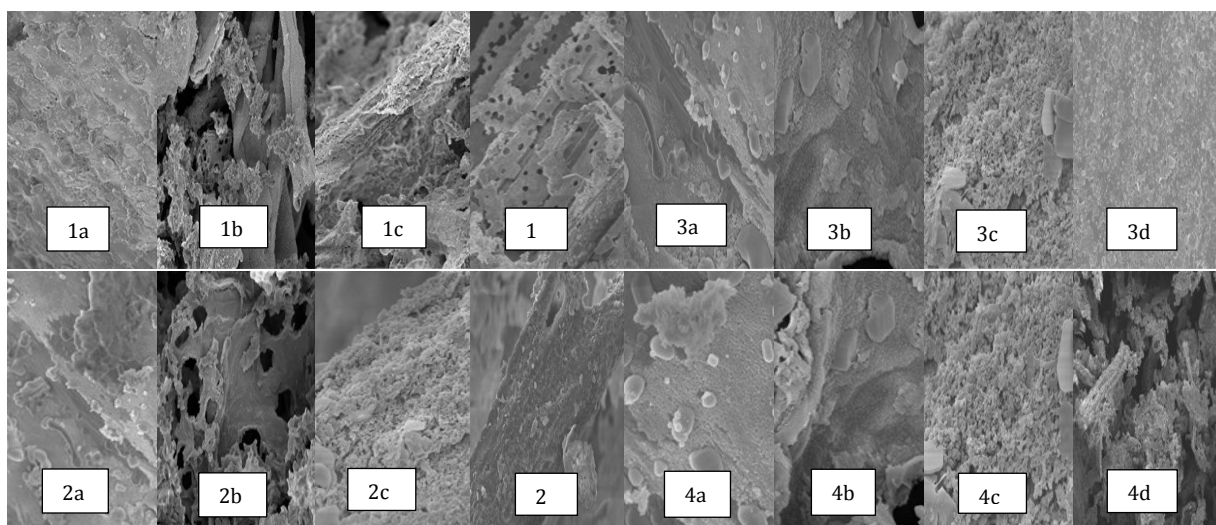


Figure 2. FESEM Mag. 1.00K and 5.00K (1a & 2a) AC_H_4a; (1b & 2b) AC_H_4b; (1c & 2c) AC_K_4; (1d & 2d) AC_OH_4; Mag. 1.00K and 5.00K (3a & 4a) AC_H_8a; (3b & 4b) AC_H_8b; (3c & 4c) AC_K_8; (3d & 4d) AC_OH_8

Conclusion

The present investigation showed that bamboo is a promising precursor to be used in the preparation of activated carbon. In the view of experiment results, the best activated carbon properties of the sample prepared from the bamboo raw material by a one-step chemical activation with base activating agent, KOH with MW-ultrasonic radiation power 300 W were obtained at the carbonization temperature of 800 °C with the highest of surface area, 1162.73 m²/g. The active surface area of the bamboo activated carbon derived in this work was considered relatively high besides being mesopores. The result proved that the potentially of MW-ultrasonic activation was rapid, efficient and feasible as a viable activation method.

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